Official Amendment
Scrial, No. – 10/038,274
Docket No. – UVD 0280 PA / 40815.324

REMARKS

Claims 1-151 were pending in the present application. Claims 1, 3, 14, 123, 126, 130, 135, 148, and 151 have been amended, claims 2, 49-122, 125, 129, 134, and 149-150 were canceled, and new claims 152-154 were added. As a result of this amendment, claims 1, 3-48, 123-24, 126-128, 130-133, 135-148, and 151-154 are pending. Reexamination and reconsideration are requested in light of the accompanying amendments and remarks.

Applicants confirm the election of Group I, claims 1-48, 123-148, and 151. Claims 49-122, and 149-150 have been canceled without prejudice pending their resubmission in a timely filed divisional application.

Applicants have updated the cross-reference information by amending the paragraph beginning on page 1, line 4.

The rejection of claims 10-11 and 14-36 under 35 U.S.C. §112, second paragraph, as being indefinite has been overcome. Claim 14 has been amended to eliminate "carbonate." Accordingly, applicants submit that the claims, as amended herein, are in compliance with 35 U.S.C. §112, second paragraph, and respectfully request that the rejection be withdrawn.

The rejection of claims 1-12, 14-15, 22, 32-33, 35, 37-42, 123-148, and 151 under 35 U.S.C. § 102(b) as being anticipated by Schapira (U.S. Patent No. 6,068,709) has been overcome. Schapira teaches a process for the phosphatization of metallic substrate comprising the use of a phosphatization bath having a pH from about 1 to 5.5 and which comprises from about 0.3 to about 25 g/l of zinc ion, from about 5 to about 50 g/l phosphate ion, and from about 0.01 to about 10 g/l of a trivalent cobalt complex represented by 1 of 2 formulas. Col. 2, lines 28-62.

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In Schapira, the trivalent cobalt complex in aqueous solution is an accelerator for the phosphating process. As discussed in "Phosphate Coatings," ASM Handbook, Vol. 5, p. 379, 1994, and "Chemical Conversion and Anodized Coatings," R.Burns et al., Protective Coatings for Metals, p. 547, 1955 (copies attached), accelerators for phosphating solutions are oxidizers, typically nitrates, nitrites, chlorates, nitroguanidine, hydrogen peroxide, etc., dissolved in the phosphating bath that increase the reaction rate with the metal substrate by climinating hydrogen formation. Schapira does not teach or suggest the use of trivalent or tetravalent cobalt in the formed coating.

Claim 1 recites that the trivalent or tetravalent cobalt/valence stabilizer complex is present in the conversion coating itself in order to provide adequate corrosion protection. Claim 1 also recites the solubility of the cobalt/valence stabilizer complex. The dependent claims further describe the physicochemical characteristics that the oxidized trivalent or tetravalent cobalt exhibits in the coating in order to provide adequate corrosion protection.

The general concept of valence stabilization of cobalt in solution was known. However, a trivalent or tetravalent cobalt/ valence stabilizer complex incorporated in a solid film for corrosion inhibition is novel.

The ability to stabilize trivalent cobalt in solution as described in Schapira does not guarantee an effective cobalt conversion coat. In the majority of cases, adding stabilizers to the bath is detrimental to the coating formed because of solubility differences. The cobalt complexes described in Schapira have high aqueous solubilities. For example, Material Safety Data Sheets for [Co(NO₂)₆]Na₃ (Aldrich Catalog No. 46,373-6) and [Co(NH₃)₆]Cl₃ (Aldrich Catalog No. 48,152-1) describe both as "very soluble in water." Although a compound that is "very soluble in water" may be an excellent choice as an accelerator for phosphating solutions, it

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would be a poor choice for a constituent compound in a conversion coating because it would be washed away quickly by corrosive media.

The high aqueous solubility of the compounds in Schapira is evidenced in the steps required to form the compounds and separate them from solution. For example, processes involving [Co(NO₂)₆]Na₃ are described in Chemical Abstracts Vol. 26, 3200, Chemical Abstracts 1628, and Chemical Abstracts Vol. 45, 4167 (copies of which are enclosed). [Co(NO₂)₆]Na₃ is used as a water-soluble precipitating agent for potassium. The sodium salt of Co(NO₂)₆ exhibits high water solubility, whereas the potassium salt is very insoluble in water. Schapira does not describe the use of the potassium salt because of its low solubility, which would make it unavailable in solution as a phosphating accelerator. The formation of [Co(NH₃)₆]Cl₃ from aqueous solution requires "salting out" with the addition of 400 ml of concentrated HCl to 1500 ml water with 230 g of [Co(NH₃)₆]Cl₃ contained therein. Inorganic Syntheses, Vol. 2, pp. 217-218. Preparation of [Co(NO₃)(NH₃)₅](NO₃)₂ from the equally high solubility [Co(NII₃)₅CO₃]NO₃ and nitric acid is detailed in Inorganic Syntheses, Vol. 4, p. 174, 1953 (copy attached). 100 ml of methanol must be added to the 25 ml of water (a standard "salting-out" procedure), followed by heating for 18 hours at 100°C to drive water off in order to produce 10 g of the desired compound. Similarly, the synthesis of [Co(NH₃)₅CO₃]NO₃ requires extensive evaporation to begin to obtain the product (1.6 I to 500 ml), Inorganic Syntheses, Vol. 6, p. 173-174 copy attached), demonstrating that even the use of carbonate as a trivalent cobalt valence stabilizer in solution does not ensure that a sparingly soluble trivalent cobalt compound will result. Synthesis of [Co(oxalate)en2]Cl, which is analogous to Schapira's [Co(oxalate)en2]NO3 also requires extensive evaporation to acquire a product. (Mori, et al. Bull. Chem. Soc. Japan 31:291-5, 1958; CA 52: 18060) (copy attached).

These examples show that there is a significant difference between stabilization in solution, where maximum solubility is desirable, and stabilization in a formed coating, where sparingly

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soluble characteristics are desired. Schapira is concerned with the presence of trivalent cobalt as an accelerator for phosphating baths. Because of this requirement, high solubility in the water bath is necessary. If the solubility of a compound in water is too low, then the compound will be unavailable in the bath to act as an accelerator. Conversely, the present invention involves the presence of a sparingly soluble compound in the formed coating. Nowhere in Schapira are the solubility requirements for the trivalent compounds discussed. Schapira's trivalent compounds are unusable as coating constituents in the present invention because of their high solubility, a necessary requirement to act as an accelerator for the phosphating bath.

The examiner stated that the term "sparingly soluble" as in claims 148 and 151 did "not distinguish over the solubility of Schapira wherein the solubility of Schapira is sufficient to form a conversion coating solution that is capable of forming a conversion coating on the substrate. The term 'sparingly' is a relative term and does not distinguish over any value of solubility." However, the specification defines the conditions of a sparingly soluble complex at p. 8, line 11, to p. 9, line 5, and p. 19, line 7 to p. 20, line 23. Schapira's highly soluble materials neither teach nor suggest this property.

According to the examiner, "the properties recited in claims 2-7, 12, and 140-143, including the solubility, the electrostatic barrier, the ion exchange property, the thickness of the coating (i.e., the coating resulting from the contact with the substrate), and the cavity containing cobalt and an additional ion are inherent properties necessarily present from the presence of the same claimed chemicals, namely the trivalent cobalt complex that has been combined with a ligar d (i.e. a valence stabilizer)." However, as explained above, the presence of a stabilized cobalt compound in solution as in Schapira is not the same as a cobalt/valence stabilizer complex within the solid corrosion-inhibiting conversion coating. Moreover, Schapira's cobalt compounds do not have the claimed solubility, as discussed. The dependent claims further describe the physicochemical characteristics that the oxidized trivalent or tetravalent cobalt

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exhibits in the coating in order to provide adequate corrosion protection, characteristics which are not inherent in Schapira's cobalt stabilized in solution.

With respect to claim 12, the cobalt compounds of Schapira do not inherently have a central cavity containing a cobalt ion and an additional ion. Nowhere in Schapira is there any type of cobalt compound which has a central cavity containing a cobalt ion and an additional ion. See p. 135, line 5-17. The examiner must provide a basis in fact and/or technical reasoning to reasonably support the determination that the allegedly inherent characteristic necessarily flows from the teachings of the applied prior art. MPEP 2112. All organic compounds and simple inorganic compounds bond directly to the cobalt ion, whether in solution or in the coating. The valence stabilizers with a central cavity are the larger inorganic valence stabilizers that can polymerize, i.e., the heteropolymetallates, such as molybdate, tungstate, vanadate, etc. Sec p. 26, line 3 to p. 28, line 16, and p. 134, line 8 to p. 135, line 20. None of Schapira's compounds have a central cavity having another metal ion in it. The central cavity only contains the trivalent cobalt ion.

With respect to claims 123-148, Schapira describes a phosphating bath, which requires the presence of a phosphate source. The phosphate source requirements are discussed in U.Evans, Metallic Corrosion and Passivity and Protection, 1948 p. 604-605, R.Burns et al., Protective Coatings for Metals, 1955, p. 545-552 (copies enclosed). The present invention does not use phosphate ions, yet provides formulations that equal or approach the performance of hexavalent chromium.

There are two general classes of conversion coats - phosphating and chromating. The present invention is not designed to replace phosphating (although it could) because phosphate baths rarely include hexavalent compounds. Rather, the present invention is designed to replace

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hexavalent chromium in chromate conversion coatings, which do not include phosphate. See Burns et al. p. 552-553.

Schapira specifically distinguishes between the phosphating process and the chromate conversion coating process.

In that connection, it is important to state that until now (see the document EP-A-0 458 020) the only known use of trivalent cobalt complexes was of their use in the surface treatments of substrates essentially based on aluminum in view of the formation on these surfaces of conversion coatings instead of carcinogenic hexavalent chrome derivatives; phosphatization treatments were not contemplated; the thus obtained conversion coatings contain aluminum oxide as major constituent, at least in volumetric percentage, and cobalt oxides CoO, Co₃O₄ and Co₂O

Col. 3, lines 1-13.

Claims 123 and 151 as amended recite "a corrosion-inhibiting conversion coating bath consisting essentially of a solvent, a precursor cobalt source, and a valence stabilizer combined to form a cobalt/valence stabilizer complex, optionally an oxidizer, optionally preparative agent, and optionally a solubility control agent." There are no phosphate ions in the bath.

Thus, claims 1-12, 14-15, 22, 32-33, 35, 37-42, 123-148, and 151 under 35 U.S.C. § 102(b) are not anticipated by Schapira.

The rejection of claims 1-12, 14-15, 123-126, 129-148, and 151 under 35 U.S.C. § 103(a) as being unpatentable over Applicant's Admitted Known Prior Art in Specification has been overcome. According to the examiner, "Applicant's Admission includes teachings of cobalt complexes having the claimed valences and valence stabilizers (e.g. inorganic or organic stabilizers) (page 5, line 23 to page 7, line 6)."

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However, contrary to the examiner's position, these references do not anticipate the claimed invention. Claims 1 and 148 recite that trivalent or tetravalent cobalt is present in the conversion coating itself in order to provide adequate corrosion protection. They also recite the solubility of the cobalt/valence stabilizer complex in the coating. The dependent claims further describe the physicochemical characteristics that the oxidized trivalent or tetravalent cobalt exhibits in the coating in order to provide adequate corrosion protection.

In contrast, as described in applicant's specification at page 5, line 23 to page 7, line 6, the references cited by the examiner are all directed to "stabilizing" the trivalent oxidation state of cebalt while it is in aqueous solution, e.g., to increase the bath life, etc. None of these references teach or suggest the functionality of trivalent or tetravalent cobalt within a solid, corresion-inhibiting conversion coating formed on a substrate metal. In fact, stabilizing trivalent cobalt in solution does not necessarily produce an effective solid, corresion-inhibiting cobalt conversion coating. In the majority of cases, adding stabilizers to the bath is actually detrimental to the coating formed due to solubility differences, as discussed above.

As noted on page 5 of the present application, the cited prior art uses additives called "bath stabilizers". These chemical species form more stable coordination bonds with trivalent cobalt cations than with divalent cobalt cations in an aqueous conversion coating solution. Specifically, carboxylates, hydroxyalkyl amines (aminoalcohols, such as triethanolamine), or nitrito complexes are described as being added to the bath to retain trivalent cobalt in solution and to stabilize concentrations during the coating process. However, unlike the present invention that is directed to maintaining oxidized cobalt (i.e., trivalent and/or tetravalent cobalt) in a solid, corrosion-inhibiting conversion coating formed on a substrate metal, the prior art bath stabilizers only treat and extend the service life of the trivalent cobalt in the conversion coating solution.

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Moreover, in contrast to the present application, "bath stabilizers" used in the manner of these specifications reduce the formation and precipitation of trivalent cobalt-containing solids during coating deposition. Bath stabilizers described in these cited references behave similar to masking agents for chemical gravimetric analysis to keep unwanted compounds from precipitating, and actually serve to starve the deposited coating of trivalent cobalt by shifting the equilibrium away from the formation of a solid, corrosion-inhibiting conversion coating containing trivalent cobalt on the metal surface, to the trivalent cobalt remaining dissolved in the coating bath (see comparative example 2 on pages 313-314). The compounds formed from the solutions described in the cited references have lower structural stability in the coating, as well as higher aqueous solubility, than if no bath stabilizer were used at all. The solubilities of the formed compounds are too high to afford long-term corrosion protection. Stabilizing the valence of the ion in aqueous solution does not suggest its application in the solid film. The idea of using valence stabilized high oxidation state cobalt incorporated into a solid film for corrosion inhibition is novel.

The examiner stated that the term "sparingly soluble" as in claims 148 and 151 did "not distinguish over the solubility admitted by applicant wherein the solubility of the composition admitted by applicant is sufficient to form a conversion coating solution that is capable of forming a conversion coating on the substrate. The term 'sparingly' is a relative term and does not distinguish over any value of solubility." However, the specification defines the conditions of a sparingly soluble complex at p.p. 8, line 11, to p. 9, line 5, and p. 19, line 7 to p. 20, line 23. The cited prior art neither teaches nor suggests this property.

The examiner also stated that "the properties recited in claims 2-7, 12, and 140-143, including the solubility, the electrostatic barrier, the ion exchange property, the thickness of the coating (i.e., the coating resulting from the contact with the substrate), and the cavity containing cobalt and an additional ion are inherent properties necessarily present from the presence of the

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same claimed chemicals, namely the trivalent cobalt complex that has been combined with a ligand (i.e. a valence stabilizer)." However, as explained above, the dependent claims further describe the physicochemical characteristics that the oxidized trivalent or tetravalent cobalt exhibits in the coating in order to provide adequate corrosion protection, characteristics which are not inherent in the cited prior art.

Therefore, claims 1-12, 14-15, 123-126, 129-148, and 151 would not have been obvious to one of ordinary skill in the art at the time the invention was made.

The rejection of claims 43-48 under 35 U.S.C. §103(a) as being unpatentable over Schapira in view of Ouyang (U.S. Patent No. 5,505,792) has been overcome. Ouyang is cited as teaching coloring and coloring additive for conversion coatings. However, Ouyang does not remedy the deficiencies of Schapira. Therefore, claims 43-48 would not have been obvious to one of ordinary skill in the art at the time the invention was made.

Applicants gratefully acknowledge the Examiner's statement that claims 13, 34, and 36 would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims. New claim 152 includes the subject matter of claims 1, 12, and 3; new claim 153 includes the subject matter of claims 1, 14, 32, and 34; and new claim 154 includes the subject matter of claims 1, 14, 35 and 36. No new matter has been added. In light of the Examiner's indication of allowable subject matter, applicants respectfully submit that these new claims are in condition for allowance.

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CONCLUSION

Applicants respectfully submit that, in view of the above amendment and remarks, the application is now in condition for allowance. Applicants respectfully request that claims 1, 3-48, 123-124, 126-128, 130-133, 135-148, and 151-154 be passed to allowance.

If the Examiner has any questions or comments regarding the present application, he is invited to contact the undersigned attorney at the telephone number indicated below.

Respectfully submitted, DINSMORE & SHOHL LLP

By 🖄

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PLP/amm Encls.

ASM Handbook®

Volume 5 Surface Engineering

Prepared under the direction of the ASM International Handbook Committee

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First printing, December 1994

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Library of Congress Cataloging-in-Publication Data

ASM International

ASM handbook.

Includes bibliographical references and indexes.

Contents: v.1. properties and selection—iron, steels, and high-performance alloys—v.2. Properties and selection—nonferrous alloys and special—purpose materials—(etc.)—v.5. Surface engineering

1, Metals --Handbooks, manuals, etc. I. ASM International, Handbook Committee. II Metals handbook. TA459,M43-1990-620.1'6-90-115 ISBN 0-87170-377-7 (v.1) SAN 204-7586

ISBN 0-87170-384-X

ASM International[®] Materials Park, OH 44073-0002

Printed in the United States of America

Phosphate Coatings

PHOSPHATE COATING is the treatment of iron, steel, galvanized steel, or aluminum with a dilute solution of prosphoric acid and other chemicals in which the surface of the metal, reacting chemically with the phosphoric acid media, is converted to an integral, mildly protective layer of insoluble crystalline phosphate. The weight and crystalline structure of the coating and the extent of pen treation of the coating into the base metal can be controlled by:

- Method of cleaning before treatment
- Use of activating rinses containing titanium and other metals or compounds
- · Method of applying the solution
- Temperature, concentration, and duration of treatment
- Modification of the chemical composition of phosphating solution

The method of applying phosphate coatings is usually determined 57 the size and shape of the article to be coated. Small items, such as mus, bolts, screws, and stampings, are coated in tumbling barrels immersed in phosphating solution, targe fabricated articles, such as refrigerator cabinets are spray coated with solution white on conveyors. Automobile bodies are sprayed with or immersed in phosphating solution. Steel sheet and strip can be passed continuously through the phosphating solution or can be sprayed.

Phosphate coating: range in thickness from tess than 3 to 50 µm (0.1 to 2 mil). Coating weight (grams per square meter of coated area), rather than creating thickness, has been adopted as the basis for expressing the amount of coating deposited

Phosphate Coatings

Three principal types of phosphate coatings are in general use: zinc iron, and manganese. A fourth type, lead phorphate, more recently introduced, is operated at ambient temperatures.

Zine phosphate coatings encompass a wide range of weights and c ystal characteristics, ranging from heavy films with coarse crystals to ultrathm microcrystalline deposits. Zine phosphate coatings vary from light to dark gray in color. Coatings are darker as the carbon content of the underlying steel increases, as the ferro is content of the coating increases, as theavy metal ions are incorporated into the phosphating solution, or as the substrate metal is

acid pickled prior to phosphating. Zinc phosphating solutions containing active oxidizers usually produce lighter-colored coatings than do solutions using milder accelerators.

Zinc phosphate coatings can be applied by spray, immersion, or a combination of the two. Coatings can be used for any of the following applications of phosphating; base for paint or oil; aid to cold forming, tube drawing, and wire drawing; increasing wear resistance; or rustproofing. Spray coatings on steel surfaces range in weight from 1.08 to $10.8 \text{ g/m}^2 \text{ (0.5} \times 10^{-3} \text{ to } 3.5 \times 10^{-2} \text{ oz/ft}^2)$; immersion coatings, from 1.61 to 43.0 g/m² (5.28 × 10^{-3} to 0.141 oz/ft²).

Iron phosphate coatings were the first to be used commercially. Early iron phosphating solutions consisted of ferrous phosphate/phosphoric acid used at temperatures near boiling and produced dark gray coatings with coarse crystals. The term from phosphate coatings refers to coatings resulting from alkali-metal phosphate solutions operated at pH in the range of 4.0 to 5.0, which produce exceedingly fine crystals. The solutions produce an amorphous coating consisting primarily of iron oxides and having an interference color range of iridescent blue to recidish-blue color.

A typical formulation for an iron phosphate bath is (Ref 1):

Component	Companition, %		
Phosphate salts	12-15		
Phosphoric and	,£ . ‡		
Molyhdate accelerator	0.25 0.50		
Detergents (anionic/nonsonic)	8-10		

Basically, then, iron phosphate formulations consist of primary phosphate salts and accelerators dissolved in a phosphoric acid solution, it is the acid that initiates the formation of a coating on a metal surface. When acid attacks the metal and begins to be consumed, solution pH at the metal surface rises slightly. This is what causes the primary phosphate salts to drop out of solution and react with the metal surface, forming a crystalline coating.

All iron phosphate conversion coatings are composed of partially neutralized phosphoric acid. But all iron phosphates are not created equal. Other ingredients, such as the specific accelerator used (Table 1), hold part of the key.

Although iron phosphate coatings are applied to steel to provide a receptive surface for the bonding of fabric, wood, and other materials,

their chief application is as a base for subsequent films of paint. Processes that produce iron phosphate coatings are also available for treatment of galvanized and aluminum surfaces. Iron phosphate coatings have excellent adherence and provide good resistance to flaking from impact or flexing when painted, Corrosion resistance, either through film or scribe undereut, is usually less than that attained with zinc phosphate. However, a good iron phosphate coating often outperforms a poor zinc phosphate coating.

Spray application of iron phosphaic coatings is most frequently used, although immersion application also is practical. The accepted range of coating weights is 0.21 to 0.86 g/m² (6.9 × 10⁻⁴ to 0.26 oz/fr²). Little benefit is derived from exceeding this range, and coatings of less than 0.21 g/m² (6.9 × 10⁻⁴ oz/fr²) are likely to be nonunform or discontinuous. Quality from phosphate coatings are routinely deposited at temperatures from 25 to 65 °C (80 to 150 °F) by either spray or immersion methods.

Manganese phosphate coatings are applied to ferrous parts (bearings, gears, and internal combustion engine parts, for example) for break-in and to prevent galling. These coatings are usually dark gray, However, because almost all manganese phosphate coatings are usually black in appearance. In some instances, a calcium-modified zinc phosphate coating can be substituted for manganese phosphate to impart break-in and antigalling properties.

Manganese phosphote coatings are applied only by immersion, requiring times ranging from 5 to 30 min. Coating weights normally vary from 5.4 to 32.3 g/m^2 (1.8 × 10⁻² to 9.83 oz/ft²), but can be greater if required. The manganese phosphate coating usually preferred is tight and fine-

Table 1 Effect of accelerators on the weight of an iron phosphate coating

		Coating weight		
Accelerator	Suchare treated	K,W ₇	114/17 × 10-1	
	Steel only	0.1140.27	0.35-0.88	
Merallic	Mixed metal loads, lerrous, zmc, and aluminum	0.22-0.38	0.71-1.24	
Oridizer	High-quality steel only	0.43-0.86	1.41-2.82	

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Table 2 Accelerators used in phosphate coating processes

	Accelerator	ru-	ative cont	menes attacks	()ptiem.		ng conditio			
Type of accelerator		٦.	у/1.	Bypl× to-	Katto		elajina.	Addition	Advantages	Limitatores
NOT:	NaN-Da, Ze(NO3)2. Ni NO205	1.3			High NOT POT	65.03	נינין ניבן		Lower shirter	Reduction of FePO ₄ increases the arm content of the anating.
KO3	NaV) ²	•	0.1 0.2	0.8-1.7	NO ₂ :NO = 1:1	inj	(a)	Continuous	Affinels, rapid processing even at low temperatures.	Corresive fumes. Highly unstable at high both temperatures. Frequent addition is required.
cio _š	Zir(ClOx)?	0.5-1				fal	(a)	Continuous	Stable in liquid concentrates. Can be used for both makeup and replemshined. Overcomes the white staming problem.	Corrosive nature of chlurase and its reduction products. High concentrations pursuan the bath. Removal of pelatinnis precipitate from the residual phosphate contagt is difficult.
H ₂ O ₂	H ₂ O ₂		0.05	0.4		ļuj	fal		Low coating weight. No harmful products. Free from stanning.	Bath control tends to be critical. Heavy shelps formation, Limited stability. Continuous addition is required.
Perheante	Sodi im perborate	•••			•••	(n)	(a)	•••	No separate neutralizer is required. Gund corrosion resistance.	Continuous addition is required Voluminous studge.
Nitroguandine	Nicrogramidine		•••			55	130		Neither the occuberator not as reduction products are correlave.	Slightly soluble, Does not control the bodding of ferrous iron in the bath. Highly expensive.

grain, rather than loose and coarse-grain. However, desired crystal size varies with service requirements, in many instances, the crystal is refired as the result of some pretreatment (certain types of cleaners and/or conditioning agents based on manganese phosphate) of the metal surface.

Manganese-iron phosphate contings are usually formed from high temperature baths from 90 to 95 °C (190 to 200 °F).

Composition of Phosphate Coating

All phosphate coatings are produced by the same type of elemical reaction: the acid bath, containing the coating chemicals, reacts with the metal to be coated, and at the interface, a thin film of solution is neutralized because of its attack on the metal. In the neutralized solution, solubility of the metal phosphates is reduced, and they precipitate from the solution as crystals. Crystals are then attracted to the surface of the metal by the normal electrostatic potential within the metal, and they are deposited on the cathodic sites.

When an acid phosphate reacts with steel, two types of iron phosphate are produced: a primary phosphate, which enters the coating; and a secondary phosphate, which enters the solution as a soluble iron compound. If this secondary ferrous phosphate were oxidized to a ferric phosphate, it would no longer be soluble and would precipitate from the bath. Cxidizing agents are incorporated to remove the soluble secondary ferrous phosphate because the ferrous phosphate inhibits coating formaticn.

Although all phosphating baths are acid in nature and to some extent attack the metal being couled, hydrogen embrittlement seldom occurs as a result of phosphating. This is primarily because all phosphating baths contain depolarizers or oxidizers that teact with the hydrogen as it is formed

and render it harmless to the metal. In some instances, however, zinc-phosphate processes, intended for use with rust-inhibiting oils for corrosion resistance or manganese-phosphate freatments, can cause hydrogen embritlement because they may contain a minimum amount of depolarizers and oxidizers. A dwell time before use or mild heating may be needed to relieve embrithement.

The acidity of phosphating baths varies, depending on the type of phosphating compound and its method of application. Immersion zinc phosphating baths operate in a pH range of 1.4 to 2.4, whereas spray zinc phosphating solutions can operate at a pH as high as 3.4, depending on the bath temperature, from phosphating baths usually operate at a pH of 3.8 to 5. Manganese phosphating baths operate in a pH range comparable to that of the immersion zinc phosphating solutions. Lead-phosphate solutions are usually more acidic than any of the others.

Zinc, iron, and manganese phosphating baths usually contain an accelerator, which can range from a mild exidant, such as nitrate, to one of the more vigorous nitrite, chlorate, peroxide, or organie sulfonic acids (Table 2). The purposes of these accelerators are to speed up the rate of coating, to oxidize ferrous iron, and to reduce crystat size. This is accomplished because of the ability of the accelerators to oxidize the hydrogen from the surface of the metal being coated. Phosphating solution can then contact the metal continuously, permitting completeness of reaction and uniformity of coverage. Accelerators have an oxidizing effect on the dissolved iron in the bath, thus extending the useful life of the solution. Some zine and iron phosphating processes rely on oxygen from the air as the accelerator. Zinc phosphating baths for aluminum usually contain complex or free fluorides to accelerate coating formation and to block the coating-inhibiting effeet of soluble aluminum.

Applications

On the basis of pounds of chemicals consumed or tons of steel treated, the greatest use of phosphate coatings is as a base for paint. Phosphate coatings are also used to provide:

- A base for oil or other rust-preventive material
- Lubricity and resistance to wear, galling, or scoring of parts moving in contact, with or without oil
- · A surface that facilitates cold forming
- Temporary or short-time resistance to mild corrosion
- A base for adhesives in plastic-metal laminations or rubber-to-metal applications

Phosphate Coatings as a Base for Paint

The useful life of any painted metal article depends mainly on the durability of the organic coating itself and the adherence of the film to the surface on which it is applied. The primary function of any protective coating of paint is to prevent corrosion of the base metal in the environment in which it is used. To accomplish this purpose, the method of preparing the metal should reduce the activity of the metal surface, so that underfilm corrosion is prevented at the interface between paint and metal.

When used as a base for paint films, phosphate coatings promote good paint adhesion, increase the resistance of the films to humidity and water soaking, and substantially retard the spread of any corrosion that may occur. A phosphate coating retards the amount of corrosion creep, because the coating is a dielectric film that insulates the active anode and cathode centers existing over the entire surface of the base metal. By insulating these areas, corrosion of the surface is arrested or at least substantially retarded.

PROTECTIVE COATINGS for Metals

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SECOND EDITION



REINHOLD PUBLISHING CORPORATION

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430 Park Avenue, New York

PAGE 52777 * RCVD AT 8/16/2004 4:11:29 PM [Eastern Daylight Time] * SVR:USPTO-EFXRF-1/1 * DNIS:8729306 * CSID:9372230724 * DURATION (mm-ss):42-58

General introduciion

American Chemical Society's Series of Chemical Monographs

The Council of the American Chemical Society, acting through its pany, Inc. (Reichold Publishing Corporation, successor) of New York. The Trustees arranged for the publication of the ACS Series of (a) Scientific and (b) Technological Monographs by the Chemical Catalog Com-New York; and the late Professor Gellart Alleman of Swarthmore College. Society and the National Research Council mutually agreed to care for these two fields of chemical progress. The American Chemical Society named as Trustees, to make the necessary arrangements of the publication of the Monographs, Charles I. Parsons, secretary of the Society, Washington, D. C.; the late John E. Teeple, then treasurer of the Society. Scientific and Technologic Monographs on chamical subjects. At the Society, chould undertake the production and publication of Critical stion with the American Chemical Society and the American Physical Tables of Chemical and Physical Constants. The American Chemical By arrangement with the Interallied Conference of Pure and Applied Chemistry, which met in Landon and Brussels in July, 1919, the American Chemical Society, was to undertake the production and publication of same time it was agreed that the National Research Council, in cooper-

Connuittee on National Policy, appointed editors (the present list of authority in their respective fields and to consider critically the manuwhom appears at the close of this sketch) to select authors of competent

scripts submitted.

the frontiers of knowielge. Furthermore, most workers in the sciences were coming to see the artificiality of the separation. The methods of three years of experience certain medifications of general policy were indicated. In the beginning there still remained from the preceding five decades a distinct though arbitrary differentiation between so-called By 1944 this differentiation was fast becoming nebulous. Research in private enterprise had grown apace and not a little of it was pursued on both groups of workers are the sance. They employ the same instra-"pure science" publications and technologic or applied science literature. The first Monograph of the Series appeared in 1921. After twenty-

Library of Congress Catalog Card Number: 56-6622

Printed in the United States of Arnemica

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6. Chemical Conversion and Anodized Coatings

Within recent years protective coatings that are neither of the metallic nor the paint type have come into prominence. They provide an insulating barrier of exceedingly low solubility between a metal and its environment through conversion of the metal surface into a corrosion-resistant.

In momentive form. The conversion consists of a chemical or electrochemical strandification of the metal surface so that the coating so formed is an integral part of the parent metal. This is in contrast to paints and metalline coatings which require no chemical combination with the basis tion with paints, enumely, and lacquers, and it is here that they find their widest applications. The protection provided is enhanced many-fold through improvement of adhesion, restriction of moisture penetration.

In through improvement of adhesion, restriction of moisture penetration and prevention of the spread of underfilm corresion. Most such firms are onserved in providing an ideal base for protective oils and waxes. In some cases conversion coatings are used without further treatments.

rates conversion variety are according treatments have attained narked commercial significance are iron and steel, and aluminum, time, and magnesium alloys. To a lesser extent cadmium, tin, copper, brass, and examless steel are finished in this manner.

and stainless steel are interest in this manner. The techniques in common use today for forming this type of protective coating fall into two general categories (1) chemical dip, spray, or brush and (2) electrolytic methods. In chemical dip reactions an oxidized form of the metal being treated or of a metal ion present in the bath is formed on the metal being treated or of a metal ion present in the bath is formed on the metal surface, the operating conditions being controlled to provide the properties required for the intended service. This tategory includes phosphating, used generally to provide a paint base on steel and zincicluronating, of value in protecting zinc and cadmium in bunild atmospheres; and chemical oxide contings, used to enhance the corrosion resistance and appearance of aluminum, steel, and copper alloy parts. Some oxidizing treatments for forming a paint base on magnesium also

fall in this entegory. Electrolytic processes, as the name implies, depend on an externally applied voltage to promote the formation of protective films in a suitable electrolyte. The foremost application of this technique is in the anodicing of aluminum alloys. Electrolytic methods are also used for several magnesium treatments.

PHOSPHATING

Phosphuting processes are used to provide protective coatings for a tion with paint, enamel, lacquer, oils, or waxes. Parts which are to be treated with oil or wax are given a thick phosphate coat. Nuts and holts are illustrative of such articles. The relatively heavy, absorbent coating retains oil for extended periods of time, excluding moisture and providing during which a thin phosphate coating is deposited. Here the coating serves to anchor the paint alm and suppress corrosion. Phospharing wide variety of iron and steel products. Zine is also phosphated in commercially significant quantities, and, to a fesser extent, aluminum alloys, cedualum, and tin. For most applications the phosphate coating is not itself sufficiently protective, but serves as an excellent base for impregnaindication. Surfaces that are to be painted are given a "fast" treatment offers the advantage of requiring only simple and relatively inexpensive equipment. Moreover, coating formation depends only on contact between the work and the buth; hence, intricate parts offer no particular problem. less, permits the use of spray application, particularly advantageous in treating bulky articles or in coating composite units after assembly, thus The development of "fast" solutions, with treatment times of a minute or avoiding damage to the coating during assembly.

Historically, phosphating is of ancient origin. There is evidence that the use of phosphates for the protection of certain iron articles was known to the Romans in the third century A.D. In 1869 it was observed" that rusting of the metallic parts of women's corects was retarded by treating them with phosphoric acid. The first important commercial application of phosphate coatings followed the proposal of Coslett in 1908 to treat iron and steel parts in a solution of phosphoric acid and ferrous sulfate in definite proportions. Later Coslett substituted zinc sulfate for the ferrous sulfate, and in 1918 it was found by the Parker Company' that

(N. Y.), 9, 207 (1911).

^{&#}x27;Jacobi, L., "Das Romerkestel Saskburg bei Homburg vor der Hohe," pp. 155, 201. 233. Homburg, 1897; Marchia O., Ind. Mescanica, 17, 817 (1835).

^{*}Tunner. R. R., Monthly Rev. Am. Bleetrophilers' Soc., 21, No. 6, 29 (1934).
* Phoenton-Muray, M., Enginsering, 85, 870 (1998); Blassett, E., Metal Ind.

C. C. C. M. d. Met. Eng., 18, 264 (1918); Eckelmann, L. E., Chem. & Met. Eng., 21, 727 (1919); Rais Moterials, 5, 488 (1922).

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the character of the coating was improved by the use of a solution of primary manganous phosphate. In this process, which came to be known as "Parkerizing," manganese dioxide was added to the bath, it being claimed that oxidation of at least some of the phosphate to the form of ferrie phosphate was essential. The time required for treatment was usually from 3 to 4 hours. While the Coslett process known as "Coslettizing" was rather widely used in England for phosphating the steel parts of bicycles and other articles, neither it nor the original Parkerizing process was suited to the needs of the rapidly expanding finishing industry.

The Parker process was improved in 1926 by the adoption of mangamese dihydrogen phosphate for the processing solution. This made possible a reduction in time of treatment. Somewhat later a further reduction in processing time to about 10 minutes was effected by incorporating a small percentage of a copper salt in the bath. This permitted the use of the process in a conveyorized or line finishing system." The increasing use of phosphate coatings as a base for paint was reflected in the name "bonderizing," given to the improved process. Still later improvements have been in the direction of time reduction and spray application.

Phosphate coatings consist of crystalline salts of the metal being treated or of metal ions added to the phosphating solution. Although the detailed compositions of phosphate baths are largely proprietary and the exact reactions complex, simple equations may be used to illustrate the formation of the phosphate coatings. A bath commonly used to coat iron or steel is made from primary zinc phosphate in aqueous solution with phosphoric acid which ultimately yields the tertiary sait, a reaction promoted by elevated temperature and agitation.

$$3Z_{\rm B} (H_2 PO_3)_2 \Rightarrow Z_{\rm B_3} (PO_4)_7 + 4H_3 PO_4$$
 (1)

The bath is so balanced that the tertiary salt is near saturation, Iron, when immersed in such a bath, is attacked by the free phosphoric acid, giving primary and secondary iron phosphate, and liberating hydrogen on cathodic areas.

$$Fe + 2H_{0}PO_{\bullet} \rightarrow Fe (H_{0}PO4)_{0} + 2H$$

$$2H \rightarrow H_{0}$$

$$(3)$$

$$(4)$$

$$F_{c} (H_{2}PO_{4})_{2} \rightarrow F_{e}HPO_{4} + H_{3}PO_{5}$$
 (4)

As the pH at the metal-solution interface is increased, the solubility products of the secondary and tertiary salts are exceeded. The over-all reaction is illustrated by:

Fe +
$$3$$
Zn (H₂PO₄)₂ \rightarrow Zn₃ (PO₄)₂ + FeHPO₄ + 3 H₃PO₄ + H₂ (5)

⁵ U.S. Palent 1,888,189.

⁴ Darney, V. M., Ind. Eng. Chem., 27, 1142 (1935).

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The crystalline layer formed is largely the insoluble tertiary zine phosphate, together with a small amount of secondary iron phosphate. Recent evidence indicates that some iron from the solution becomes an integral part of the coating during the process. The proportion of iron to zine is dependent on the method of application, solution composition, temperature, and time of processing.

The solution composition is critical in that too much free acid results only in pickling of the steel surface,* while too little promotes sludge formation. This second effect becomes increasingly troublesome as the concentration of ferrous ions in the bath builds up with time. Although it is felt that some iron phosphate is beneficial to the adherence and protective qualities of the coating, excessive amounts are detrimental to corrosion resistance.

A second salt frequently used in phosphating baths is manganese phosphate. The reactions are analogous to those for zine except that the initial reaction (Eq. 1) yields the secondary as well as tertiary salt.

The coating reaction illustrated above is an extremely slow one, owing largely to cathodic polarization. Accelerating agents are therefore added to the bath to reduce treatment time. Oxidizing agents, such as nitrates. nibrites, or clorates, behave as depolarizers in oxidizing the hydrogen formed (Eq. 3). Other accelerators for increasing the reaction rate include copper salts, organic compounds such as nitrobenzene, bases such as aniline, toluidene, pyridene, and quinoline. Provided the conditions are so controlled that a fine grain size is obtained, there is no appreciable sacrifice in protective value associated with the use of accelerators. Through the use of accelerators, reaction times have been reduced to less than a minute for paint-base contings. It has thus become possible to apply phosphate coatings by spray techniques as one step in the conveyor production of large items such as automobile budies, fenders, etc. The chart in Figure 16-1 summarizes the phosphate treatments in current use together with the basis metals treated and methods and purposes of application.

As in any coating process, the success of the operation depends on proper preparatory surface treatment. Mechanical surface cleaning, such as emery treatment, or blasting with sand, grit, or shot, is employed to remove scale and rost. Degressing is accomplished with trichlorethylene or in an alkaline bath. Phosphate coatings are particularly sensitive to

Eisler, S. L., and Chamberlain, P. G., Metal Finishing, 50, No. 6, 113 (1952).

^{*} Phosphoric acid cleaning, as distinguished from a true phosphating treatment, employs a high-acid bath primarily to remove surface dirt and grease, leaving a slight a riace each to promote mechanical bonding with paint, together with a very thin presphate residue. To insure proper cleaning, the acid concentration must be so high that a substantial crystallino phosphate film cannot be developed.

PROTECTIVE COATINGS FOR METALS

grease, hence parts should not be handled with bare hands after degreasing-

The protective qualities of phosphate coatings are largely a function of the crystalline character of the film, protection increasing with de-

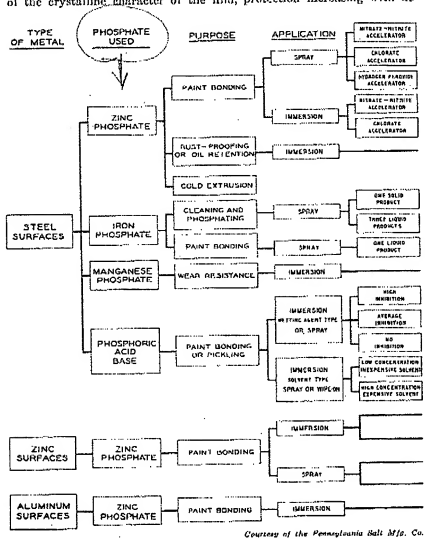


FIGURE 16-1. Summary of types and uses of phosphate coatings.

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creasing grain size. In general, refinement of the crystalline structure is obtained with increasing temperature. By using suitable addition agents, the temperature can be reduced with no adverse effect on grain size. Crystal structure is also affected by physical treatment of the surface prior to phosphating. The "wiping" effect, whereby the surface is mechanically wiped prior to treatment, results in refined crystal structure and improved corrosion resistance. This same effect can be approached cherrically in a solution containing 1 per cent sodium phosphate and 0.01 per cent titanium as the anhydrous phosphate salt. The work is precipped in this solution; then, without rinsing, into the phospate bath. This treatment is applied to both iron and zine, but primarily to the

Applying phosphate films to ferrous surfaces is an accepted method of reducing wear and galling on mating parts. Properly lubricated, the coating provides myriads of oil reservoirs, effective in reducing friction. Furthermore, the coating is deformed slightly under pressure, thus reducing maximum applied stress. Heavy phosphate films for this purpose are ordinarily formed from "slow" baths of zinc or manganese phosphate at a temperature between 90 and 100°C (200–210°F). Reaction times range from 10 to 60 minutes, to produce films of 1000 to 4000 mg per sq. 4. There is little dimensional change associated with these coatings: it normally is of the order of 0.005 mm (0.0002 inch) for a heavy coating. Of the many applications of this type, many are found in the automotive industry. Piston rings, rocker arms, camshafts, valves, tappets, and cylinder liners are frequently phosphated to increase life.

The use of sine phosphate coatings to reduce friction in wire, tube, and deep drawing applications was suggested about twenty years ago, but was not commercially developed until the recent war. Coatings of 150 to 3000 mg per sq/ft are used, the thicker coatings being specified where deformation is light or moderate. For severe forming operations the thickness is limited to preclude injury to the coating. Lubricated coatings of this nature, by markedly reducing friction, reduce wear on dies, permit faster forming operations, and conserve power. No adverse effects on physical properties result from phosphating. Hardness, tensile strength, and temper are unimpaired.

Although the foregoing friction-reducing uses of phosphate coatings are commercially important, the phosphating treatment is known and

^{*} Reeves, M., Metal Ind. (London), 78, 7, (1951).

¹ Jernstedt, G., Trans. Electrochem. Soc., 83, 351 (1943).

¹⁰ Roosa, M. B., Imbrication Eng., 6, 117 (1950).

¹³ Ayres, R. F., Materials & Methods, 34, No. 4, 100 (1951).

^{: *} Holden, H. A., Sheet Motal Ind., 26, 123, (1949).

PROTECTIVE COATINGS FOR METALS

used primarily as a paint base for increasing corrosion resistance. The perous nature of the film provides good mechanical anchorage for paint, enamel, or lacquer; the phosphate layer prevents moisture which penetrates the paint film from reaching the basis metal and stiffes the spread of corrosion beneath the paint film. These coatings are formed in "fast" zinc or manganese phosphate solutions at temperatures ranging from room temperature to 210°F. Treatment times generally are from 30 seconds to 5 minutes. Coatings range from 100 to 600 mg per sq ft. The risk of corrosion from phosphating solution retained on the treated part varies with the type of solution. Plain metal phosphate-phosphoric acid types are not particularly corresive, but some of these containing accelerating agents can be quite harmful." An after treatment in a chromate bath is frequently used to provide added protection. A chromic acid rinse, with acid concentrations up to 0.5 per cent, often proves beneficial, maximum performance being associated with 0.1 to 0.2 per cent. Concentrations greater than 0.5 per cent are injurious because of the dissolving action of the rinsing bath.

Phosphate coatings, being essentially insoluble in most service environments, are protective insofar as they are continuous; continuity, hence, protection, increases with thickness. The corrosion resistance of phosphated but unpainted steel is compared with that of untreated steel in Table 16-1.14

Table 16-1. Corrosion of Priosphated Steel (SAF-1020) Compared with That of Untrated Steel

COMPARED WITH LEAVE		Phosphatod (mg/cm²)
Test 744 hours in salt log	Untreated (mc/em²) 34.08 + 20.39	3.22 1.60

The extent to which phosphate coatings aid in the retention of paint on

steel panels is shown in Table 16-2.13 In addition to increasing adhesion and retention of paints, phosphate coatings also tend to stille under film corrosion. A recent six-year test illustrated the protection provided by two spray coats of baked synthetic enamed applied over steel samples treated in three different ways: (1) solvent vapor degreasing, (2) sandblasting, and (3) coating with phosphate. Whereas underfilm and general corresion were evident on the first

in Clarke, S. G., and Longburst, E. E., J. Iron Steel Inst. (London), 170, 15 (1952). A Darsey, V. M., and Cavanaugh, W. R., Parker Rust Proof Co., Trans. Electro-

¹⁰ Darsey, V. M., Parker Rust Proof Co., "Phosphate Coatings," Reprinted with chem Soc., 91, 351 (1947). permission from "Corrosion Handbook," edited by H. H. Uhlig, p. 867, New York, John Wiley & Sons, Inc., 1948.

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two surfaces, the phosphate coating improved paint protection and retarded underfilm attack.16

The affinity of phosphate coatings for oil and wax is exploited in rustproofing many articles. The adhesion and abrasion resistance of the coating is such that threaded articles, such as nuts, bolts, and screws, can be

Table 10-2. Refertion of Paint by Untreated and Phosphate-Coared Steel

	Printer Wt, Retained (oz/fl²)	Increase in Retainten Over Untreated Street (%)
Steel untrented	0.139	
Steel-phosphate coated Roller application	0.177	27.3
Steel-phosphate coated By spray	0.200	43.9
Steel-phosphate conted By immersion dipping	0.242	74.0

so treated. During World Wars I and II small arms were rustproofed by phosphating, then treating with chromic acid and a rust-preventive oil. The corrosion protection obtained is indicated in Table 16-3, is in which are listed salt-spray requirements for military equipment finished in this manner.

PAGLE 16-3. SAUT SCHAY RESISTANCE OF PROSPITATED STEEL BEFORE AND AFTER OLLING

Class	Conting Wt. (mg/ft*)	Salt Spray Resists Before	After
A	1000-4000	1	24
Ts	1000-4000	2	36

Besides iron and steel, other metals, notably zine and aluminum, are phosphate treated on a large scale. Zine surfaces which are painted with no intermediate treatment provide inadequate service due primarily to the reaction of zine with the paint vehicle. Zine soaps are formed, destroying the bond between the paint and the zine. Phosphating is one treatment used to avoid this difficulty, and also to increase the mechanical anchorage of the paint. Cadmium is treated in the same way. The baths are of zine phosphate in which the work is dipped for about 2 minutes, forming a film of 75 to 200 mg per sq ft. Phosphating is recommended frequently for zine- or cadmium-plated steel that is to be painted. Zine-flash bonderizing consists in applying a thin electrodeposit of zine which is given a phosphating treatment, then painted. This composite coating provides a higher degree of corrosion protection to iron or steel than does normal phosphating and painting. The zine thickness is

h

¹⁶ Gentieu, N. P., Machine Design, 24, No. 2, 141 (1952).

¹⁷ Douty, A., Plating, 38, 1031 (1951).

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552 usually 0.0006 to 0.0015 mm (0.000025 to 0.00006 inch) for indoor and 0.0025 to 0.0035 mm (0.00010 to 0.00015 inch) for outdoor exposure.

For treating tin a special technique is available called "Protecta Tin" in which an oxide and phosphate complex are formed on the tin, imparting an increased resistance to corrosion and to staining by sulfur products.16 Processing is carried out by immersion for less than 30 seconds at 85°C in a solution containing 9 pounds trisodium phosphate (anhydrous), 8 pounds sodium dichromate, 20 pounds sodium hydroxide, 3 pounds wetting agent, and water to make 121 gallons.19 The treatment does not affect the appearance of the metal nor does it interfere with subsequent lacquering or lithographic printing. This process is used to increase the shelf-life of canned foods both by retarding rusting of the outside of the can during storage in damp conditions, and by preventing the unsightly blackening of can interiors resulting from sulfur originating in meat, fish, soup, and many vegetables.

CHROMATE COATINGS

Chromate conversion coatings were widely used during World War II for protection of zine- and cadmium-plated parts in tropical service. Since that time they have undergone marked commercial development, and are now firmly established in the metal finishing field. They are used purely for protection purposes without additional finishing treatment, or they may be finished with lacquer or paint to combine protection with product appearance. Their use has been extended to the protection of zinc-base die castings and aluminum.

Zinc, when exposed to conditions that give rise to moisture condensation, develops a bulky white corrosion product. This most readily occurs in highly humid conditions such as prevail in the tropics or the Gulf Coast where the daily temperature variation causes intermittent condensation or dew formation. Storage of zinc or zinc-coated parts in unheated warehouses during periods of high humidity or the packaging of such products in boxes made of unseasoned wood or damp paper stock has caused corrosion of this "white bloom" type. In addition to detracting from the appearance of the part, these corrosion products can impair the operation of certain types of equipment, such as relays and switches. It is for such applications that chromate coatings are largely employed to retard the corrosion of zine and cadmium plated parts and zinc-base

die castings. Applications of chromate coatings are predominantly by immersion

¹⁸ Kerr, R., J. Soc. Chem. Ind., 65, 101 (1946).

¹⁰ Product Finishing (London), 3, No. 12, 58 (1950).

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processes, although electrolytic methods are used to some extent in the ai-craft industry. The bath compositions are largely proprietary but all centain two basic constituents: hexavalent chromium ions and a mineral acid. Some also contain one or more organic acids. On immersion, the zinc or cadmium article is attacked by the mineral acid with an attendant rise in pII of the solution next to the metal. At the same time some hexavalent chromium is reduced to the trivalent state. At a critical pH the trivalent chromium and some hexavalent chromium are coprecipitated on the metal surface.

One of the oldest and still popular chromating processes is known as the "Cronak" process concerning which details have been published. The bath contains chiefly sodium dichromate in fairly high concentrations, slightly acidified with sulfuric acid. Analytical studies indicate the film formed from this bath to be a basic chromium chromate of the general formula Cr.O. CrO. xII.O. A typical film, dried at 110°C (230°F), has been analyzed as follows:

Hexavelent chromium	8.63% 28.2%		
Trivalent chromium	3.27%		
Sulfate			
Zinc	2.12%		
Sodium	0,32%		
Water	19.3%		

The thickness is about 0.0005 mm (0.00002 inch).

Chromate films are generally noncrystalline, nonporous, and gel-like. They are quite susceptible to damage from abrasion immediately following film formation; hence should be aged 12 to 24 hours, before being subjected to normal shop handling. Electrolytically applied coatings are not subject to this limitation.

Chromate films which provide maximum protection contain both trivaent and hexavalent chromium.23 Protection is provided in two ways. The comporous nature of the film physically excludes, to a large extent, moisare from the metal. At discontinuities in the film, the hexavalent chromium, being slightly soluble, exerts its well-known inhibiting action. Experiments show that the formation of white corrosion product is inhibited so long as a minimum amount of hexavalent chromium remains in the film. The useful life of a chromated part thus depends on the rate at which hexavalent chromium is leached from the film, i.e., on the degree of wetting of the part. The protection provided by a Cronak coating in stagnant water is illustrated in Table 16-4.21

MONAME

Registered trade mark of New Jersey Zinc Company.

²⁵ U.S. Patent 2.035,380 (Mar. 24, 1930) E. J. Wilhelm (to New Jersey Zine Co.).

²¹ Anderson, E. A., Proc. Am. Electroplaters' Soc., 30, 6 (1943).

²² Ostrander, C. W., Plating, 38, 1033 (1951).

· Chemical Abstracts

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ZnI4 m. 132 3"; (E2S);ZnI4 m. 149"; (E2MeS);ZnI4 m. 173-4"; (E2PrS);ZnI4 m. ALEREO HOFFMAN

Composition of complex metal eyanide radicals. Complex Ni eyanide radical (MASAKI) 2. Synthesis and x-ray investigation of NiCr₂O₄ (Holderson) 2.

LAPHMA, HISNEI: Recherches sur les aluminates de calcium et sur leurs combinaisons avec le chlorure et le sulfate de calcium. Paris: Vuibert. 68 pp. F. 16,

7-ANALYTICAL CHEMISTRY

W. T. HALL

An improved method of quantitative spectrographic analysis. C. C. Nitteniz and C. W. Standin. Ind. Eng. Chem., Anal. Ed. 4, 182 85(1932). The method employed in a modification of the methods of Carlach and Schweitzer and N. (C. A. 23, 1587). In the improved method an addal, element is added to the sample in a const. known and, for comparison purposes. The use of a recording microphotometer is now justified and considerable time is saved. Four requirements are fisted for satisfactory comparison of spectral lines. App. and procedure are described, and spectra and microphotometric records are shown. Concrete examples are given and the errors are analyzed. The deviation from the mean of 4 detms, is 1 part in from 10 to 20. H. A. S. Micro-alkalimetry and addimetry. S. K. Concov. Bull. inst. recherches biol. Perm 7, No. 7/8, 427-40(German summary) 440-4(1931). The solo, to be investigated is titrated to the intermediate color of a 2-color indicator, with not stronger than 0.01 N reagents from a membrane microborat. Formulas are given for caleg. the decrease in acidity or alky. pn values are taken from tables.

Thermometric titration. Takayuki Somiya. J. Soc. Chem. Ind. 51, 135-407 (1932); cf. C. A. 21, 1425, 3030; 23, 1840, 2907, 4907, 4908; 24, 3380, 3915.—The necessary app. is shown and numerous applications of the method are described.

Use of heaviding acetate as indicator in molybdo-manganimetry; application to 1587). In the improved method an addul, element is added to the sample in a const.

Use of benziding acetate as indicator in molybdo-manganimetry; application to the electrolytic determination of a few hundredths of a milligram of copper. R. Curlinthe electrolytic determination of a few hundredths of a milligram of copper. R. COULUS-mer AND L. THIYOLLE. Compt. rend. soc. bink. 108, 30, 2(1021).—Fontés und Thivolle (C. A. 17, 1199) developed a method for detg. Cu which depended upon the fact that when Cu is plurged into a phosphomolybilate reagent (boil 40 g. of NII, molybilate, 60 ec. of NaOH solm. 4, 1.36, and 100 ec. of water until no more NII, is evolved, cool, add 200 ec. of water and 200 ec. of 11, PO₄, d. 1.30, boil 15 min, cool and dilt to 1 t.) the Cu dissolves, forming a blue soln, which becomes colorless when titrated with KMIOa. To overcome the difficulty in detg. the end point, it is now recommended to use benzidine acctate soln, as indicator. Dissolve 1 g. of benzidine in 10 ec. of glacial AcOH, hoil and dilt, to 100 ec. After washing an electrolytic Cu denosit, plunge the electrode into enough of the phosphomolybdic reagent to cover it (3 ec. in a small electrode into enough of the phosphomolybdic reagent to cover it (3 cc. in a small tube with a Pt wire cathode), allow the color to develop for a few min, while using the cathode as a stirrer and titrate with dil, KMnO, after adding the indicator. W. T. H.

A neutral buffered standard, for hydrogen-lon work and accurate thrations, which can be prepared in one minute. ROUGH, WILLIAMS AND CARL M. LYMAN. J. Am. Chem. Soc. 54, 1911-2(1932).—NH₃OAe solus. through wide limits of conco. have a pn of almost exactly 7.00.

W. T. H. pn of almost exactly 7.00.

Pu or almost exactly 1.001.

New reagent for the detection of hydroxy acids. C. H. Linuxalli. Eal. assoc. braxil. pharm. 12, No. 10, 23(1931).—Hydroxy acids can be detected by the may of the following reagent: FeCls (10%) 32.4 cc., KCNS (10%) 58.2 cc., H₂O to 100 cc. The reagent turns yellow on the addn. of a neutral solut of hydroxy acid. AcOU and exactly acid also give the same color. Addn. of I drop of HNO₂ brings back the original red color with hydroxy acids. With AcOH and oxalic acids the color does not return.

Preparation of sodium cobaltinitrite as potassium reagent. E. Rupp and A. Postendore. Apoth. Zig. 47, 282-3(1932). To 5 g. of powd. $Co(NO_3)_2$ in 2.1 g. AcOH and 1 g. H_2O add a luke warm soin, of IO g. NaNO₄ in II g. H_2O , remove the NaO₄ by a 30-min, current of aspirated air through the mixt., allow to stand 15–30 min. pass through a filter to remove any turbidity and mix in a pure-lain mortar with 20 g. of pure, freshly dehydrated Na₂SO₄, then place in a vacuum desiceator for a day. Triturate the golden yellow cryst, mass and preserve in amber glass. W. O. E.

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Chemical Abstracts.

then placed in H₂O until they sink and finally immersed for 12-24 hrs. in the SoCl, sol, mentioned above. After washing in water, they are put successively into alc., xylene, and parafliu. The shreds can now be investigated under the microscope after the removal of the paraflin, or else treated previously according to Pappenheim; the Hg is visible in little globules. By animal experiments, the author has determined that on keeping the cadaver, the reaction gradually disappears during the decomp, process, probably owing to the formation of HgS; in general, the SoCl, reagent does not give good results after 6 days.

J. W. Bain.

The Precipitation of Fischer's Sait Ordinarily Supposed to be Potassium Cobaltinitrite. L. L. Dekontack. Bull. smc, belg. chim., 23, 200-2.—For the precipitation of K, Dek. recommends a sol, of 50 g. cryst. Co(NO₂)₂ dissolved in 1 l. H₂O acidified with egree. HNO₂ (d. 1.2). This sol, is added to a sol, of 300 g. cryst. NaNO₂ dissolved in 1 l. of 11₂O. By the use of this mixture. E can be found in amounts as small as 11.10,000, while with the HOAe sol, formerly recommended, the limit was 11.2000. Dek. also reports that better results are obtained in separating Ni and Co by using a sol, containing 35 g. NaNO₂ and 22 g. KNO₂ in 100 cc. H₂O in place of the 40% KNO₂ sol, formerly employed. The reason is obvious since com. KNO₂ is far more impure than com. NaNO₂.

The System: Water, Pyridine and Sodium Carbonate. A Study of Solubility and the Carves of Saturation. H. Lamboscu. Bull. soc. balg. chim., 23, 179-200.

I. K. PRE

MINERALOGICAL AND GEOLOGICAL CHEMISTRY.

ROGER C. WELLS.

Recent Methods of Rock Analysis. P. G. W. BAYLY. Melbourne. Austral. Assoc. Adv. Sci. Meeting of Jan., 1907. Reprint from author, to pp.—This is in part an argument in favor of uniformity of method in rock analysis, based on the work of the chemists of the U. S. Geol. Survey, in order to make possible comparisons of the greatest real value, and a plen for an organized classification of the igneous rocks of Australia on the lines adopted by the American Survey. A few slight modifications of the methods of Hillebrand as laid down in Bull. 176 are worked out and compared as to results with those obtained by the original methods. At least two of the modifications had already been adopted by Hillebrand in Bull. 30s of the U. S. Geol. Survey before the above paper was tend.

W. F. Hillebrand,

Obsidianites—Their Origin from a Chemical Standpoint. H. S. Semanas. Proc. Roy. Soc. Victoria, 21 [2], 423-43—Six recorded and three new analyses of obsidianites are compared by the American system of classification of igneous rocks. They show compositions rarely met with among terrestrial rocks, most of them falling in the group almerose, from which type the reat do not greatly diverge. There is nothing in common between their comp, and that of rocks from places that have been named as possible sources. There is an apparent provincial distribution, and, if so, their cosmic origin is the only possible one. Billitonites are shown to be genetically connected, with the obsidianites, and for moldavites, which have few representatives among terrestrial rocks, the argument in favor of a cosmic origin is strengthened.

W. F. Halbandand.

Does Thorium Exist as Thorium Silicate in Monazite? Outo Kress and F. J. Metzorr, Columbia Univ. J. Am. Soc., 31, 640-52, figs. (Kress, Diss., 17 pp.).—Thirty distinct crystals from North Carolius and Norway were analyzed. Large crystals were broken into fragments and each fragment analyzed separately, whereby

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exchanged at once. Fo³⁺ and FeY⁻, both 0.0125 M, at pH from 1.5 to 5.0 showed an immediate partial and very slow subsequent exchange. Temp. = 25°. Tracers were Co[∞], Fe[∞], and Ni[∞]. Cf. C.A. 42, 1523a. P. E. B.

New approaches to the study of the reaction between a potassium and cobaltinitrite long. Angel del Campo, Vicente Boissier, and Angel Floyos (Univ. Madrid). Rev. real acad. cienc. exact. fix. y nat Madrid 34, 352-69(1940).—KaCo(NO₂)₈ was pptd. with stirring by adding DeKoninek's reagent (C.A. 3, 1623, (1909)) to a known soln. of KCl, filtering, drying in a current of air at 100°, and weighing in a tared filter enneible. The wt. of ppt. is less when the soln. is stirred during pptn.; hence adsorption takes place. On long standing, with the ppt. in contact with an excess of b NaCo(NO₂)₈, the wt. increases considerably; Na probably substitutes for K in the ppt., as evidenced by qual, analysis. The ppt. is probably a mixt. of K₁(Co(NO₂)₈], kfo and K₁Na (Co(NO₂)₈], but the K is nevertheless pptd. quant. K, pptd. with Li₁(Co(NO₂)₈], can be filtered and washed more easily than when pptd, with the Na reagent, and comes down as well-formed cubic crystals corresponding closely to the formula K₂(Co(NO₂)₈] when an excess of reagent is employed; the wt. of the ppt. is almost independent of whether or not the soln, is agirated during pptn., in contrast to the large changes observed with the Na reagent. The anion of the K the soln, is agitated during pptm, in contrast to the large changes observed with the Na reagent. The anion of the K R. L. Wolke soln, has no effect.

aoln, has no effect.

Chemistry of therium in aqueous solutions. 1. Organic and inorganic complexes, R. A. Day, Jr., and R. W.—Stoughton (Oak Ridge Natl. Lab., Oak Ridge, Tenn.). J. Am. Chem. Soc. 72, 5602-7(1950).—Equil. consts. for the formation of complex ions of Th with the anions of HF. HIO3, Cl.CCOOH, HBrO3, Cl.CHCOOH, 11NO3, HCl. and CICH-COOH in aq. soln. with a H-ion cough. of 0.5 M and d an ionic strength of 0.5 were obtained by detg. the distribution ratios between the aq. soln. and benzene contg. thenoyl-triducroacetone (1) an a chebuting agent. The coosta. for the 1:1 complexes decrease in the order of the above listing, going from 4.3 × 10⁴ for ThF²⁺ to 1.33. Higher complexing corresponding to Th/anion ratios of 2 and 3 was

found for the first five anions in the list; the corresponding

lound for the first five anions in the list; the corresponding consts, are presented. Evidence was found for the existence of double complexes of the type Th(NO₂)F²⁺ and Th(NO₂)F₂ in mixts, of HNO₂ and HF. Acetic acid in the aq. phase increases the extractability of the chelated product formed between Th and (1).

Sequestering properties of salicylic acid. Complexes of salfosalicylic acid. Giovanni Mannelli. Ann. chim. applicata 38, 594-601(1948).—Fe is sepd. from Tl by treating a soln, of the salfates with NH, salicylate (1) and (NH₂)CO₂, followed by pptn. of Fe with H₂S. Addn. of aq. NH₂ and holling then gives TiO₂. Fe is sepd. from Mi, with I, the Mn being pptd, as phosphate, and similarly Fe from Tl, the boiling then gives 1103. For is sopil from Mil, wan I, the Mn being pottl, as phosphate, and similarly Fo from II, the II being pottl, as chromate in presence of I. Na sulfo-salicylate and Be(O11), give Na, Be sulfosalicylate, Culta-O₁₈S₂Na,Be; similarly are formed Na,UO₂ sulfosalicylate, Culta-O₁₈S₂Na,Be; similarly are formed Na,UO₂ sulfosalicylate, Culta-O₂₈S₂N Ni

HaOnSaKtNI.

Electrochemical study of complexes of the silver ion and Electrochemical study of complexes of the silver for and amino dicarboxylle acids. Suzame Valladas-Dubois. Compt. rend. 231, 1299-1300(1950).—Aspartic acid and glutamic acid, like glycine, at high pH form complex anions of about the same stability, corresponding to 2 mols, of the anino acid to one of AgNO₃. Earl S. McColley amino acid to one of AgNOs.

Atsenates—oxidation of assenious acid (Takahashi) 4. Kinetics and mechanism of solid-phase reactions—in oxide films on pure I/e (Cadbransen) 2. Heavy-metal aluminides (Rühenback) 9.

Taurins, A.: Neorganiska Kimija, Würtenberg: A. Brokana N. R. Dovana. 1948. 126 pp. Wells, A. F.: Structural Inorganic Chemistry. New York: Oxford Univ. Press. 1951. 500 pp. 37. Experimentelle Einführung in die Anorganische Chemic. 44th ed. Edited by W. Klemm and W. Fischer. Berlin: W. de Canyter. 1949. 204 pp. DM 7.80. Reviewed in Angew. Chem. 63, 55(1951).

7-ANALYTICAL CHEMISTRY

History of analytical chemistry in Russia. K. B. Yatsimirskii. Uspekhi Khim. 18, 623-8(1949).

Recent developments in inorganic analytical chemistry. J. C. Geertsma. S. African Ind. Chemist 2, 31-3(1948).—A description is given of (1) the use of the Walden Ag reductor and internal indicators in the detin. of Re with f K,Cr₂O₇, (2) detin. of C in presence of large autra. of S, e.g., and in pyrites concentrates, with MnO₂ on pumies granules to absorb the SO₂ and SO₂, and (3) detin. of CO₂—in solid nucterials.

B. A.

materials. Analytical methods for determining the concentration of ionically active substances. II. Schwerdtner (Chemnitz, Ger.). Chem. Tech. (Berlin) 2, 361-4(1950). A review. Paul W. Howerton

Qualitative analysis without hydrogen sulfide. X. Reduction method. Toshiyasa Kiba (Kanazawa Univ.) J. Chem. Soc. Japan, Pure Chem. Sect., 70, 145-7(1949); cl. C.A. 45, 2813c.—Metallic Zn can be used as a group reagent to sep. ions from acidic soln. Systematic procedures of group sepn. and identification of ions are given in tables. tables.

Expansion of the percentages, \$\pmu 20%; \$\pm 20%;

Quantitative analysis of powders by infrared spectro-photometry. Georges Pirlot (Univ. Liège). Bull. soc. chim. Bolges 59, 327-51(1950); cf. C.A. 43, 7366h. The compensated extinction coeff, method, detd, with an interval standard, is applied to the analysis of powders when Beer's law does not hold, and when there is overlapping hetween the absorptions of the components of the mixt. Results are given for the analysis of mixts, of cinchonidine, cinchoniue,

quinidine and quinine; and of lencine, norlencine, and iso-II. N. Dyer lencine.

seneme.

Spectrochemical analysis of a solution on graphite. M. van Doorselaer. J. Escelahott, and J. Gillis (Univ. Ghent. Beig.). Congr. graupement numee. mithod gnal. spectrograph. produits met. 12, 51-7(1919).—Factors which affected bronze analysis according to the method of Rivas (cf. C.A. 32, 1601) were studied. When adsorption of the soln. on graphite took place at a temp. above 100°, the sensitivity of spectral lines was at a max, and their relative intensities were const. An increase in the conen. of the soln, cave a of spectral lines was at a max, and their relative intensities were const. An increase in the conen, of the soln, gave a greater sensitivity, except in the case of some elements such as Al, but did not influence the relative intensities. The effect of the vol. of the soln, on the electrode was negligible. A change in the anion ratio of a soln, produced changes in the spectra because of differences in the volatility, as well as adsorption and absorption characteristics of various saits and their complexes. The presence of colloidal Sn gave abnormally high results.

M. A. Rinehart normally high results.

normally high results. M. A. Rinehart
Amperometric titration of suifhydryl groups: microgram
analysis. Sheldon Rosenberg, J. C. Perrone, and Paul L.
Kirk (Univ. of California Med. School, Berkeley). Anal.
Chem. 22, 1186-7(1950).—The argentometric procedure of
Kolthoff and Harric (C.A. 44, 76a) is modified. A vibrating Pt electrode is used as a combination electrode-stirrer,
and the fitrating soln, is added from a horizontal microburet. For y annes, of cysteine-HCl and glurathione the
error was less than 5% in all cases. Titration of denatured
horizon sarum albumin should approx. 0.3% sulfhydryl (as
cysteine): this value was essentially independent of the
nature of the denaturing agent.
T. H. Dunkelberger

eysteine); this vatue was essentiary independent of the nature of the denaturing agent.

T. H. Dunkelberger Utilization of ion exchangers in analytical chemistry. XVI. Gunnar Cabrielson and Olof Samuelson (Chalmers Tek. Högskola, Göteboug, Sweden). Swensk Kom. Tid. 62, 214-20(1950)(In English); cf. C.-4. 44, 6759f.—Aldehydes may be retained quantitatively by an ion exchanger in the bisulfite form. The break-through eneves have the same that the state of the same than a detained for inner mode, showing that the shape as those obtained for inorg, acids, showing that the

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120

A yield of 35 g. of the hydrous 3-1-hydrate is advanted from The process of settling in the later water, centrifuged, and dried at 110° in an electric oven. stages is very slow and much time may be saved by the use The precipitate is finally washed with 124, nortions of 100 g. of FeCl.:611.O. of a centriluge.

Analysis

dissolving a weighed portion in warm, concentrated nitric dissolved in ammonia, reprecipitated with dilute miric acid. After evaporation to remove excess acid and cillution procedure. Typical analytical results give an Fe:Cl ratio with water, the chloride is precipitated as silver eliforide, acid, and weighed in the usual manner. The ironitt in The amount of chleride in the sample is determined by ammonia and is weighed as Fe_sO, according to the standard the combined filtrates is desermined by precipitating with of 40:1.

References

- 2. Betoeson and Hartong: Induganic Stweeters, 1, 185 (1939).
 - Wriser and Millionn; J. Am. Clem. Soc., 67, 235 (1935).
 - 3. Weiter and Millioan: Chem. Ress., 25, 1 (1934).

59. HEXAMMINECOBALT(III) SALTS*

Stemmer of Janes Bierrun and James P. McReynheds Checked by Alpred L. Opperability R. W. Pabay Hexaconinecobalt (III) salts may be prepared by any of three methods that depend on oxidation of cobalt(II) ion

by Dk. McReywolds, who was impressed by the simplicity of Dr. Bjornan's *Procedures for preparing most of these salts were originally submitted method. Shortly after the manuscript for Vol. II had been sent to the publisher, similar procedures for three of these salts were submitted by D. Bjerran. Insamuch as the method was developed by Dr. Bjerran and the editors have taken the liberty of combining the two sets of directions wer conditions prevented his submitting the procedures at an earlier deco, and presenting them as a joint effort of the over men.

| University of Contendagen, Consectagen, Denmark.

The Onio State University, Columbus, Oking depensed.

Chiversity of Minole, Urbana, III.

HEXANDHNECOBALPANI, SALTS

(2) exidation with an agent such as hydrogen perexide, in ammoniacal solution: (1) air oxidation, with formation mine by heating with aqueous ammonia under pressure; 173 13 chlorite solutions;2 (3) oxidation in the presence of a catalyst that allows equilibrium between the pentammine and hexammine ions to be established at room temperature and atmospheric pressure. These compounds may also of the pentaromine ion, which is converted to the hexanlodine, * potussium permangamate, * lead dioxide, * or hypobe prepared indirectly from other hexamminecobalt (III) Salts, 3, 13

ion 12.13 or decolorizing charcoal 11 is used as a catalyst. charcoal is the catalyst is simple, gives high yields of pure product, and is not time-consuming. A high concentration of ammonium salt is sufficient to stabilize the hexanthe equilibrium. Air is used as oxidant except when the In the best of the catalytic methods diamninesilver The method devised by J. Bjerram's in which decolonizing minecobalt(III) ion, and the carbon serves only to establish cobalt(II) compound is slightly soluble in the ammoniacal solution, as in the preparation of [Co(NH₃)₅]Br₃, for which hydrogen peroxide is preferable.

Procedure

A. HEXAMMINECOBALT(III) CHLORIDE

4[Co(NH₅), JCl₃ + 2H₂O $4\text{CoCl}_2 + 4\text{NH}_1\text{Cl} + 20\text{NH}_3 + 0_2 \rightarrow$

chloride 6-hydrate and 160 g. (3 mols) of ammonium chloride are added to 200 ml. of water. The mixture is activated decolorizing chargoal* and 500 ml. of concen-Two hundred and forty grams (1 mol) of cobalt(II) shaken until most of the salts are dissolved. Then 4 g, of trated ammonia are added. Air is hubbled vigorously * In order to obtain a rapid reaction, care must be exercised in the choice Charecal from the center of a package has been found to he much more effective than that from the top. of this cetubyet.

SESTIFIC SENTIFIES

sh brose insually about 4 hr.).* The air inlet tube is of fairly large hove (10 mm.) to prevent elogging with the prebrough the mixture until the red solution becomes yellowcipicated hexauminecobalt (III) salt.

be used to give the entire mixture an acid reaction. The and is filtered hot. The hexamennecobalt(III) chloride is chloric acid and slowly cooling to 0°. The precipitate is The erystals and carbon are filtered on a Büchner funnel mixture is heated on a hot plate to effect complete solution precipitated by adding 400 rd. of concentrated hydrofiltered, washed first with 60 per cent and then with 95 per cent alcohol, and dried at 50 to 100°, i Vield 230 g. (85 laydrockloric acid in 1500 ml. of water; sufficient acid should and then added to a solution of 15 to 30 ml. of concentrated per cent). And. Culvd. for [Co(NH₁)] [Cl₄: Co, 22.0; Cl₄ 39.8; NH, 38.2. Found: Co, 22.0; Cl, 39.8; NH, 38.1.

B. HEXAMMINECOBALT(III) NITRATE

1. 4Co(NO₃)₂ + 4NH₄NO₅ + 20NH₅ + O₅ +

4[Co(NH;)6(NO;); + 2H;O

 $(Co(NH_2)_0]Cl_0 + 3HNO_0 + [Co(NH_2)_1](NO_2)_1 + 3HCl_1$

Method 1. Seventy-three grams of coball nitrate 6hydrate (0.25 mol) is dissolved in 100 ml. of water, and, in ammonia (about 2.5 mols of N.H.,) are added. The solution is exidized by air as specified in Procedure A. The salt swashed with a little ice-took water and is dissolved on a steam bath in 1300 to 1500 ml. of water containing enough uitric acid to give the mixture an acid reaction. After the order mentioned, 80 g. (1 mol) of armrouium nitrate, 2 g. of activated charcoal, and 180 ml. of concentrated

* Ours should be taken that the building is not too vigorous, as this removes part of the armannia from the solution. If this happens, the solu-tion stays red instead of becaming yellow. Attiough more aumonia may he added, the yield will be degreesed somewhat

! When the calt is dried at a Ligher temperature, it sometimes rurns green, The original color is reserved by redissolving and reprecipitading with nytho-

chleric seid.

HENAMAGINECOBALITATIO SALTS

A THE PARTY OF THE

The presidented and is marked with water and alcohal and dried at 100°. Yield 77 to 79 g. (88 to 90 per cent). Anal. emoval of the carbon by filtration, the solution is treated with 200 mi, of concentrated mitric unid and left to cool Caled. for [Co(NH₂),[NO₄]; Co, 17.0; NH₃, 29.4. Co, 16.9; NH, 29.4.

and dried at 100? Yield 103 g. (88 per cent). A solution of these crystals gives only a slight turbidity with silver cobalt. II) chloride (Procedure A) is dissolved in 1500 ml. of The garbon is removed and the salt precipitated with 450 ml. of concentrated mitric acid. The crystals are washed with two 200-ral, portions of alcohol to remove excess acid The precipitate from the exidation of water containing 15 to 30 ml. of concentrated nitric acid. Method 2. eitrete.

C. HEXAMMINECOBALT(III) BROMIDE

2[Co(NH₃)], Br₃ + 2H₂O. $C_0CO_1 + 2HBr - C_0Br_2 + H_2O + CO_2$ 2CoBr₂ + 2NH, Br + 10NH, + H, O₂ -

When the vigorous effertoscence has ceased, the into the desired product. To complete the equilibrium adjustment, the mixture is heated for 5 min. on a steam mixture is filtered, washed with a little cold water, and peroxide (0.40 mol) is added slowly while the solution is main part of the cobalt(II) precipitate has been converted bath and then left standing for 1/4 hr. The saft-carlion Twenty-four grams (0.20 mol) of cobalt carbonate* is added slowly to 100 ml, of 45 per cent hydrobromic neid (0.50 mol). To the solution, 2 g. of activated charappears is disregarded, and 40 ml. of 30 per cent hydrogen reated with 900 to 1000 nd. of water containing sufficient coal and 120 ml. of concentrated annuous (1.6 mols of NHs) are added. The precipitate of cobalt(II) add that stirred.

* A cokalt determination must be made on the collect carlumnto used if on accurate check on the yield of the complex saft is desired.

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grinding procedure is repeated. The residue is again collected on a filter and washed with cold water until the filtrate is no longer orange, but instead is faintly greenish. The salt is finally washed with methanol and ether and dried at 50°. Yield, 7.8 g. (86%). Anal. Caled. for [Co(Nff₃)₅-I](NO₃)₂: NH₂, 21.8. Found: NH₂, 21.8.

D. NITROPENTAMMINECOBALT(III) NITRATE

 $\begin{array}{l} [\mathrm{Co}(\mathrm{NH_3})_3\mathrm{CO_3}]\mathrm{NO_3} + 2\mathrm{HNO_3} + \mathrm{NaNO_2} \rightarrow \\ [\mathrm{Co}(\mathrm{NH_3})_3\mathrm{NO_3}](\mathrm{NO_3})_2 + \mathrm{NaNO_3} + \mathrm{CO_2} + \mathrm{H_2O} \end{array}$

Ten grams of carbonatopentamminecobalt(III) nitrate (0.036 mol) is suspended in 25 ml, of water, and 10 g, of sodium nitrite (0.14 mol) and 10 ml, of colorless nitric acid (1:4 concentrated acid and water) are added. The mixture is stirred for 15 minutes at room temperature, and then 200 ml, of methanol is added to the slurry. The precipitate is collected on a filter, washed with methanol and other, and dried at 50°. Yield, 10 g. (88%). Anal. Calcd. for [Co(NH₂)₂NO₂I(NO₃)₂: NII₃, 27.1. Found: NH₃, 27.1.

E. NITRATOPENTAMMINECOBALT(III) NITRATE®

 $\begin{array}{c} [\mathrm{Co}(\mathrm{NH_3})_{5}\mathrm{CO_3}]\mathrm{NO_3} + 2\mathrm{HNO_3} \rightarrow [\mathrm{Co}(\mathrm{NH_2})_{5}\mathrm{H_2O}](\mathrm{NO_3})_{3} \\ & + \mathrm{CO_2} + \mathrm{H_2O} \\ [\mathrm{Co}(\mathrm{NH_3})_{5}\mathrm{H_2O}](\mathrm{NO_3})_{3} \rightarrow [\mathrm{Co}(\mathrm{NH_3})_{5}\mathrm{NO_3}](\mathrm{NO_3})_{2} + \mathrm{H_2O} \end{array}$

Ten grams of carbonatopentamminecobalt(III) nitrate (0.036 mol) is suspended in 25 ml. of water, and 20 ml. of colorless nitric acid (1:1 concentrated acid and water) is added with stirring. When the evolution of carbon dioxide has stopped (10 minutes), 100 ml. of methanol is added, the aquopentammine cobalt(III) nitrate is collected on a filter, and washed with alcohol and ether. This salt is heated at 100° for 18 hours (until 1 mol of water is lost), yielding 10 g. (83%) of the desired material. Anal. Calcd. for [Co(NH₃)_x-NO₃](NO₃)₂: NH₃, 25.8. Found: NH₄, 25.6.

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CARBONATOTETRAMMINECOBALT(III) NITRATE

safety trap and into a beaker of water where the acid is titrated with 2 N sodium hydroxide solution.

The mixture is stirred at a rapid rate and beated for 30 minutes to 126°, whereupon the black slurry thickens. Only approximately 1 ml, of the sodium hydroxide is consuncd in this initial heating. The reaction is then allowed to proceed rapidly at 128 to 139° for 2 hours. In the course of the reaction the mixture becomes light tan and appreciably less viscous, and the theoretical amount of sodium hydroxide is required to neutralize the hydrogen chloride which is evolved. Very little additional hydrogen chloride is liberated during another hour of heating.

The solid is filtered, washed repeatedly with chloroform or benzene, and dried in vacuo; the yield is 99%. Anal. Calcd. for FeCl₂: Fe, 44.06. Found: Fe, 43.89.

References

- P. Kovacic and N. O. Brack: J. Am. Chem. Soc., 76, 5491 (1954).
- 2. N. O. Brace: U.S. patent 2,719,074 (1955).

55. CARBONATOTETRAMMINECOBALT(III) NITRATE

 $2\text{Co}(\text{NO}_3)_2 + 6\text{NIJ}_2(aq_2) + 2(\text{NIJ}_3)_2\text{CO}_3 + \frac{1}{2}\text{O}_2 \rightarrow$ $2(C_0(NH_0)_4CO_2|NO_3 + 2NH_4NO_4 + 11_2O_4)$

Scemetred by G. Schlessinger* CHUCKED BY JOHN W. SIMMONE, I GERHARDT JABE, I AND MARK M. CHEMBERLAINT

The first salts of this series were prepared by Vortmann' and later extensively investigated by Jørgensen, who, however, gave only semiquantitative synthetic preparations.

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INORGANIC SYNTHESES

The method given below, which is a more exact procedure than given hitherto (and which may be significantly shortened by the use of hydrogen peroxide instead of air as oxidizing agent) may also be used, virtually without change, for the preparation of the obloride, bromide, iodide, sulfate, selenate, and oxalate of the carbonatotetranmine series simply by using an equivalent amount of the corresponding cobalt(II) salt as starting material.

Procedure

One hundred grams of cobalt(II) nitrate 6-hydrate (0.344 mol) is dissolved in 100 ml. of warm water and added to a mixture of 200 g. of ammonium carbonate (2.08 mols) in 1 L of water and 500 ml, of concentrated ammonia. The resulting liquid is oxidized by sucking air through the solution contained in two 1-1, suction flasks fitted with stoppers and glass air-inlet tubes reaching almost to the bottoms of the vessels.* After 21 hours, when the oxidation is complete, the solution is evaporated on the steam bath to a volume of 500 ml. Any cobalt(III) oxide is filtered off while hot, and further evaporation to 350 ml. is carried out. During the course of the evaporation 50 g, of solid ammonium carbonate should be added in 5-g, portions at regular intervals.† The solution is next cooled in ice, filtered by suction, and the crystals pressed well dry. The crystals may be washed with 75 ml. of alcohol. The filtrate is evaporated down to 100 ml., with the addition of four 5-g. portions of ammonium carbonate, and more product is isolated as above. The first batch of product, 48 to 50 g., is analytically pure, but the second may be contaminated with a trace of carbonatopentamminecobalt(III) nitrate, which is

A timesaving but more contly method of oxidation is to add 250 ml, of 3% hydrogen peroxide slowly to the well-stirred cobalt(II) ammine mixture. After standing 10 minutes the solution is evaporated as below. The yield is the same.

[†] The regular addition of the ammonium carbonate is emphasized.

Chemical Abstracts

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the unsym, compd. reacts more slowly. The trisulforates hydrolyze rapidly, giving SO₂=*, SO₄=*, So₄=*, and a little tetrasulfonate; azedisulfonate is an intermediate. Tetrasulfonate hydrolyzes only slightly. R. II. Jaquith

The stability of metal tetraethylenepentamine complexes. Charles N. Reilley and J. H. Holloway (Univ. of N. Caro-lina, Chapel Hill). J. Am. Chem. Soc. 80, 2017 (190958).— From potential-pH diagrams the stability consts. (pK) of the From potential-pH diagrams the stability consts. (pK) of the following metal ion complexes of tetracthylenepentamine (I) were detd.: Hg(II), 27.7; Cn(II), 22.9; Ni(II), 17.8; Zn, 15.4; Cd, 14.0; Pb(II), 10.41; Mn(II), 7.0; Bi(III), Ca. Mg, Sr, Ba, Al, and La, all negligible. pK values of 2–5 for I were found to be 4.4, 8.2; 9.2, and 10.9, resp., at 25° and with an ionic strength of 0.10. Analytical applicabilities for I complexes are discussed. Gerald S. Golden

Arsenic(III) exycbloride, a new type of polymer bond. E. Thilo and P. Flögel (Akad. Wissenselmfum, Berliu-Adlershof). Angew. Chem. 69, 754(1957)(in German). ActO₄ dissolves in AsCla with formation of an anorphous, hard polymer which has the structure Cl-AsOI(Cl)AsOI(**

Polytide I. March. Rudolph J. Marcus AsCly.

The synthesis of metal complexes. III. Synthesis of ethylenediaminecarbonato, ammineoxalato, and ethylene-callamineoxalato series of cobalt(III) complexes. Motolichi Mori, Muraji Shibata, Eishio Kyuno, and Keji Hoshiyama (Univ. Kanazawa). Bull. Chem. Soc. Japan 31, 291-5 (1958); cf. C.A. St. 56 Sh.—Three series of Co(III) conplexes were systematically synthesized from the green solu-of potassium tricarbonatocobaltate(III) (I), by means of of potassium tricarbonatocobaltate(III) (I), by means of successive substitution of carbonate by other groups: blue K[Co(en)(CO₂)₂], H₂O (II); violet K[Co(en)(CO₂)₃], H₂O (III); [Co(en)₂]Ch, H₃O (VI); [Co(en)₂]Ch, H₃O (VI); blue-K[Co(NH₂)₂(C₂O₃)₃], H₃O (VII); [Co(NH₂)₂Co₃Cl (VIII); K₃[Co-(C₂O₃)₃], H₃O (IX); blue K[Co(en)(C₂O₄)₂], H₂O (X); violet K[Co(en)(C₂O₄)₃], H₂O (X); violet K[Co(en)(C₂O₄)₃], H₂O (XI); and [Co(en)₃(C₃O₄)₃] Cl, H₃O (XII). Of these, the blue and the violet varieties are probably to be regarded as new complexes, stereoisometic with each other. Through all the procedures a green cold soln, of I was used as starring material. I was propel from with each order. Through an the procedures a green charson, of I was need as starting material. I was propel from 20 g. KHCO₆ 10 g. CoCl₂ 61₂O₅ and 5 ml. 30% H₂O₂ by the method described in the previous paper. To I was added 5 g. ethylenediamine extinuate (XIII), prept. by passing CO₂ into a cold aq. soln, of 70% ethylenediamine (XIII). passing Cer into a cold all some of 10% employments (XIV). The mixt, was placed in an ice bath for 30 min, until the green color changed to blue, and 20 ml. of abs. EOH was added. Impure crystals sept, and were tiltered off and rejected; 50 ml. addul. ETOH was added to the filtrate, and the solu, was kept at 0° for some time. Crystals of II were collected and purified by disadving them in a min and of sold II of our additing the EUOL. tals of H were collected and partial by dissolving them in a min, and, of cold H₂O and adding abs. EtOH; 4 g, of pure H were obtained. To I was added 5 g, XIII as before and the mixt, was kept 20 min, at room temp. A violet solu, was obtained; 20 ml. EtOH was added and the solu, was stored overnight in the refrigerator. The resulting ppt. was recrystd. 3 times as before with cold H₂O and abs. EtOH to yield 6 g, III. 1V was prepd, by adding 10 ml. 70% XIV to I, heating the mixt, on a water bath until the solu become reddish yellow. After cooling, 11 g, of crude soln, become reddish yellow. After cooling, 11 g, of crude product was collected and recrystal, from H₂O. V was prepal by adding 15 ml. 70% XIV, 3 g, KCl, and I g, active C to I. The mixt, was heated 30 min, on a water was bath until the color changed to brownish vellow, after which C was filtered off. The filtrate was exapt, to half the initial vol., cooled, and the ppt, washed with EtOB and dried. The yield was 12 g. V. I contg. 10 g. powd. (NII₄)₂C₃O₄ was warned to 70°; when the solit, became bluish, it was cooled in an ice bath to stop further reaction and 50 ml. EtOH was added. The sole, was allowed to stand 1 hr, and the resulting crude product was collected and signal are and the resulting crade product was conscitued and reppted, twice from the cold soin, by addin of EaOH. The final ppt., which was washed with EtOH and Et₂O in turn, yielded 7 g. VI. The starting materials, the same as for VI, remained on a water bath until the color turned blue and finally violet; then 80 ml. EtOH was added and the solution to overnight in a refrigerator. The violet ppt, was collected, dissolved in small ant. of cold H₂O, and the com-The properties of calcium silicate hydrogen was twice reported by addn. of ELOH. After the product was washed with ELOH and EL₂O, it yielded 7 g. to Berkovich, D. M. Khelker, O. I. Gracheva VII. Powd. (NIL₂)C₂O₂ (5 g.) and 30 ml. coned. NH₂OH were added to I. 2 g. NH₂Cl was added, and the mixt, was warned at 80° until the soln, became pink. The soln, was warned at 80° until the soln, became pink. The soln was solded, and the mixt was warned at 80° until the soln, became pink. The soln was solded in ml. ElOH was added to put, all other salts of the soln, became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The soln was added to put, all other salts of the soln became pink. The properties of calcium silicate hydrache was added to put and the mixt, was added to put and the mixt was added to put and the properties of calcium silicate hydrache was added to put and the properties of calcium silicate hydrache was added to put and the mixt was added to put and the properties of the properties of the properties of the propertie

which were removed by filtration, and 20 ml, addal. BOH was added; the yield was 6 g, VIII. To a cold soin, of I was added 16 g, powd, (CO-H), until a bhigh green soin, was obtained. The soin, was kept at 60° as long as CO₂ was obtained. The solu, was kept at 60° as long as CO₂ was evolved and then was cooled, acidified with dil. AcOH, and 50 ml. E(OH was added. The yield was 12 g. IX, which was purified as usual. K₂C₂O₁ (5 g.) and 5 g. ethelenediamine oxidate, preprint by neutralizing 15 g. 70% XIV with 23 g. powd. (CO₂H)₂ in the cold, were pound into a cold solu, of 1. The mixt, was kept about 1 hr. at room temp, until the color became bluids violet, after which it was cooled, treated with a track of ECOM to compare forms with with a small ant, of EtOH to remove foreign salts, and the filtrate was further treated with EtOH to ppt. 3 g. X. The ppt. was purified by washing with EtOH and Et₂O. XI ppt, was purified by washing with EtOri and Cigo. At (7 g.) was synthesized in the same way as X, except that the nixt, was heated 1-hr-instead of bring left at room temp. A mixt, of h g. (CO₂H)₅, 2 g. KCl, and 16 ml, 70% XIV was added to I and the mixt, eyapd, on a water bath until a crust formed on the surface. The solu, was cooled and the crystals were filtered and washed with EtOH and Et2O, yielding 10 g. XII. R. E. Dunbar

The study of the Raman spectra of the system hilrosyl sulfuric acid-sulfuric acid water with a view of dentrating highly concentrated sulfuric acid. Arthur Simon and Horst Richter (Tech. Rochschule, Dresden, Ger.). J. prakt. Chem. [4], S. 68-76(1957).—Raman spectra of vacuum distd. H₂SQ, were recorded with a 3-prism Zeiss spectrograph having a dispersion of 8A./mm. at 4038 A. and 12A./mm. at 4046 A. HNOSO, was prepd, from 84O₅ and HsSO₄. The N₂O₅ content of the nitroso acid was detd, with a Lunge nitrometer and U2SO4 was also detd, by the Lunge method. In the production of 14-80, from appearing nitropo acids mean which cannot be denitrated with SO₂. With the aid of Ramon spectra the conditions neces-sary for denitrating highly cound, 14,804 can be established, and the reactions between the nitroso group and SO₂ can be clarified. The NO i frequency in the Raman spectrum decreases as the concus, of water and N₂O₄ increase. The decrease being smaller the lower the conen, of H₂SO₁. It is shown that the depression is caused by HSO₁. H₂SO₁ undecides (mostly assord, by H-bridges) shield the polarizing effect of the HSO," on the NO bion and thereby change its bonding condition. Attempts to denitrate show that the bonding condition of the NO+ ions, characterized by the position of the Raman frequency, is decisive for the denitrability of nite so acids. George Meister

The complex formed from cobalt hydrocarbonyl and butadiene. Haus B. Jourssen, Robert I. Stearns, Jouko Kenttanian, Donald W. Moore, and A. Greenville Whittaker (Tolane Univ., New Orleans, La.). J. Am. Chem. Soc. 80, 2580-7(1958).—The complex formed between butadiene and Co hydrocarbonyl in the absence of Oxo conditions was prepel, and the stereochem, configuration was studied. preptl, and the stereothem, configuration was studied. When K cobaltearbonylate was allowed to react 12 hrs. with AcCill and liquid buladiene in a high-pressure vessel, a red-brown liquid was obtained, b. 23-5° at less than 1, mm., and laving the compn. Co(CO)₃C₁H₁. It is diamagnetic and contains no acidic H. The infrared and direct violet absorption curves indicate the disappearance of the conjugated dioletin structure and the appearance of 4 structure resolution of the configuration contains the configuration of the conf structure producing absorption characteristics similar to those of eis monoblefins. The sharp peak at 703 cm. bettielt in the Co hydrocarbonyl has been assigned to the B vibrations, also disappears. Felix Saunders

The interaction of ammonium and polassium hexabromoosmate(IV) and liquid ammonia at 25". II. Ammonia osmate(1V) and liquid aminoma at 25°. It. Alimnoma soluble products. George W. Watt and Lauri Vaska (Unive of Texas. Austin). J. Floorg. & Nuclear Chem. 6, 246-51 (1988); cf. C.d., 52, 8819g., 12642h.—Reaction of K.O.Bra and liquid NII₂ at 8-25° gives NII₃-3d. p-nitridubis-[dibromotetra numineosmium(IV)] brounds. This was isolated as the 3-hydrate and converted to p-nitridubis-[dibromotetra numineosmium] mium(IV)]triindute tribydrate. When (NH4):OsBre accompanied by NH Br is allowed to react similarly, hexammineosminm(III) bromide is produced by reaction of NH₄Br and accompanying hexammineosminm(III) hexag bromessmate(111). Jack J. Bulloff

The properties of calcium silicate hydrates. T. M. Berkovich, D. M. Khelker, O. I. Gracheva, L. S. Zeving and N. I. Kupreeva. Doklady Akad. Nank S.S.S.R. 120, 853-6(1958). Ca silicate hydrates C.S.H. (C = CaO S = SiO₂, and H = H-O) were synthesized. X-ray powder diagrams, electron micrographs, thermobalance, differential

METALLIC CORROSION PASSIVITY AND PROTECTION

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PROTECTION BY CHEMICAL AND

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generally considered to have value when applied to metal-work previously freed from scale, but it appears extremely doubtful whether they afford any means of escape from the necessity for descaling. All the evidence obtained by the Iron and Steel Corrosion Committee's tests at Derby, and by the tests conducted by Mayne at Cambridge, suggests that the baths mentioned above provide no substitute for pickling or shot-blasting.

Attempts have been made at Cambridge 1 to precipitate an inhibitive substance within the small amount of adherent rust which remains on steel which has been exposed to the weather and then wire-brushed; this can be accomplished by means of a process of double-spraying. Two sprays are arranged so that the solutions contained in them impinge and mix as they converge in droplets on the surface of the metal. For instance, one spray may contain zine sulphate and the other potassium chromate, so as to precipitate zine chromate, an inhibitive agent, in the interstices of the adherent rust. Numerous other pairs of solutions have been tried and Thornhill's tests show one of the best combinations to be lead nitrate and sodium phosphate. After the double-spraying, paint was applied in the ordinary way and the specimens exposed for 23 years at Cambridge. It was found that double-spraying was valueless for a freshly ground surface, but on such a surface, painting alone gives good protection, and chemical pre-treatment is not necessary. On the other hand, when applied to a surface which had been weathered for such a time as to reach the very unfavourable condition mentioned above, double-spraying greatly improved the performance of the paint. Doubtless the precipitated inhibitor did not adhere to a bright rust-free ground surface, but did remain in the interstices of the rust just at the places where it was required. This system of pretreatment has possibilities, but it does not prevent entirely the failure of paint applied to a partly weathered surface. Experiments are now being carried out by Mayne on different lines.

Production of Phosphate Layers and Similar Coatings on Iron and Steel. Phosphate processes (p. 597) have been widely adopted for the preparation of the relatively thin steel-work used in the enamelled or painted portions of automobiles or cycles. Often the bath is based on manganese dihydrogen phosphate along with free phosphoric acid; the analogous zine compound may also be present. Iron phosphates tend to accumulate in old baths, but this is probably undesirable, owing to the tendency of iron phosphate films to oxidize. At one time, long periods of treatment were necessary, but recently much more rapidly acting baths have been introduced; times as short as 7 seconds are now possible. Copper salts are valuable accelerators, but nitrates, nitrites or chlorates are also used; sometimes an organic substance (e.g. a nitro-compound) may be chosen to speed up the cathodic reaction; bases such as aniline, toluidine, pyridine or quinoline are stated to have a similar effect, whilst one process makes use of reducing agents such as bisulphites. Bary a finds that if the

- ¹ R. S. Thornhill and U. R. Evans, J. Soc. Chem. Ind., 1939, 58, 13.
- 2 J. Bary, J. Latte Corr., 1938, p. 431C.

rapid baths are so constituted as to render the grain size of the deposit very fine, they give almost as good results after painting as the slow baths.

Numerous other modifications of procedure have been proposed. Some authorities advocate the presence of sodium chromate (0.5 g./l.) in the water used for rinsing after the phosphate bath.

The process is varied according as the film is intended to form the sole protective layer, or merely to act as a basis for enamel or paint. In the latter case, porosity is probably an advantage, so that the paint may soak in. Frolably the main reasons for the good adhesion of paint are three. (1) The mill-scale and rust which would otherwise in many cases have been left below the paint are removed in a preliminary operation, before the articles enter the phosphate bath; (2) the phosphate coat appreciably reduces the danger of the local formation below the paint of bulky rust and of alkali, which are frequently causes of detachment; (3) the paint or enamel flows into the interstices between the phosphate crystals, so that the paint coat is anchored to the metal.¹

The phosphate coats are usually grey; a black finish can be obtained by impregnation with a dye, followed by oiling or greasing. There are many varieties of the phosphate treatment.² An early commerical process received the mone of Coslettizing, whilst two treatments much used to-day are known as Parkerizing and Bonderizing. Parkerizing is used for nuts, bolts, tools, and it tricate mechanisms in typewriters, cameras, and automatic machines; here the coat is scaled only with oil. Bonderizing is intended for use where a paint, counsel or lacquer, is to be applied. It is largely employed in the automobile, cycle, refrigerator and typewriter industries, also for telephone and switchboard equipment, and to some extent for taps and similar fittings

Speaking generally, phosphate treatment has proved its value as a base for lacquer or enamel. Tests by Kolke, however, show that for resisting hydrochloric or sulphuric acids, it is better to apply the enamel or varnish direct to the untreated metal; specimens on which it has been applied to a phosphate-treated basis failed promaturely, apparently owing to solvent action of the acid on the phosphate. Similar tests 4 conducted in 3% sodium chloride solution under conditions of partial immersion favoured specimens painted over a phosphate-treated surface, which, with every paint combination tested, gave results superior to the corresponding specimen painted on a ground not treated in the phosphate bath. A set of photographs published by Marshall 5 illustrate the superior resistance to salt-spray of steel treated by the Bonderizing process before pointing.

- 1 H. Sutton, J. Sci. Inst., 1943, 20, 87.
- R. C. Duvies and S. J. Scouse, J. Electrodep. Soc., 1943, 19, 13.
 H. Chaso. Met. Alloys, 1942, 15, 420.
 E. Rackwitz, Korr. Met., 1934, 10, 58.
 A Burkhardt and G. Sachs, Korrosionstagung, 1932, 2, 47.
 L. Schuster, Stald n. Eisen, 1940, 60, 786.
 G. Roesner, L. Schuster and R. Krause, Korr. Met., 1941, 17, 174.
 H. Krause, "Metallfärbung," 1937, pp. 129, 151 (Springer). Some cause: of defects are discussed by O. Macchia, Korr. Met., 1936, 12, 211.
 - ⁹ F. Kolke, Farb. Zeit., 1931, 36, 2235.
 - 4.7. Kolke, Parb. Zeit., 1934, 39, 331.
 - ⁴ F. C. B. Marshall, J. Oil Colour Chem. Assoc., 1939, 22, 242.

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